L Number	Hits	Search Text	DB	Time stamp
10	45594	chlorophenyl	USPAT;	2001/09/24 15:24
			US-PGPUB;	•
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			DERWENT;	
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			IBM TDB	
24	41560	disulfide	USPAT;	2001/09/24 15:24
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			DERWENT;	
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			EPO; JPO;	
			DERWENT;	
	55	aldrithiol	IBM TDB USPAT;	2001/09/24 15:18
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			EPO; JPO;	
			DERWENT;	
			IBM TDB	
_	16057	retrovir\$4	USPAT;	2001/09/24 10:32
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-	17	aldrithiol and retrovir\$4	USPAT;	2001/09/24 10:33
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	1005		IBM TDB	2001/09/24 10:34
-	1037	zinc adj finger\$1	USPAT;	2001/09/24 10:34
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-	62	karlstrom.in.	USPAT;	2001/09/24 10:56
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			EPO; JPO;	
	i		DERWENT;	
_	5223	hiv and protease	IBM TDB USPAT;	2001/09/24 10:57
-	3223	into and procease	US-PGPUB;	2001/05/24 10.5/
			EPO; JPO;	
			DERWENT;	
			IBM TDB	
_	4	karlstrom.in. and (hiv and protease)	USPAT;	2001/09/24 10:59
	•	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	US-PGPUB;	
	1		EPO; JPO;	
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			DERWENT;	

Search History 9/24/01 3:40:01 PM C:\APPS\EAST\workspaces\09431607.wsp

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FILE 'MEDLINE' ENTERED AT 09:28:14 ON 24 SEP 2001
L1
           6164 S ZINC(W)FINGER
L2
         103154 S HIV
            253 S L1 AND L2
L3
          23679 S DISULFIDE?
L4
            18 S L3 AND L4
L5
           3043 S MALEIMIDE?
L6
            0 S L3 AND L6
L7
           1977 S HYDRAZID?
L8
             0 S L3 AND L8
L9
             0 S ALPH (W) HALOGENAT? KETONE?
L10
             3 S HALOGENAT? KETONE?
L11
             0 S L2 AND L11
L12
L13
             0 S 2127-03-9/CRN
     FILE 'REGISTRY' ENTERED AT 10:18:43 ON 24 SEP 2001
L14
              1 S 2127-03-9/RN
                SET NOTICE 1 DISPLAY
                SET NOTICE LOGIN DISPLAY
     FILE 'REGISTRY' ENTERED AT 10:19:24 ON 24 SEP 2001
                SET TERMSET E#
                DEL SEL Y
                SEL L14 1 RN
              1 S E1/RN
L15
                SET TERMSET LOGIN
     FILE 'MEDLINE' ENTERED AT 10:19:29 ON 24 SEP 2001
             67 S L15
L16
             39 S L16 AND PY<=1994
L17
               E RICE WILLIAM G/AU
               E RICE W G/AU
L18
             51 S E3
             0 S L18 AND L16
L19
             9 S L18 AND L4
L20
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L24 ANSWER 2 OF 4

MEDLINE

ACCESSION NUMBER:

91288502

DOCUMENT NUMBER:

MEDLINE 91288502 PubMed ID: 2062837

TITLE:

Copper inhibits the protease from human

immunodeficiency virus 1 by both cysteine-dependent and

cysteine-independent mechanisms.

AUTHOR:

Karlstrom A R; Levine R L

CORPORATE SOURCE:

Laboratory of Biochemistry, National Heart, Lung, and

Blood

Institute, National Institutes of Health, Bethesda, MD

20892.

SOURCE:

PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES OF AMERICA, (1991 Jul 1) 88 (13) 5552-6.

Journal code: PV3; 7505876. ISSN: 0027-8424.

PUB. COUNTRY:

United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

Priority Journals

FILE SEGMENT: ENTRY MONTH:

199108

ENTRY DATE:

Entered STN: 19910825

Last Updated on STN: 20000303

Entered Medline: 19910802

AΒ The protease of the human immunodeficiency virus is essential for replication of the virus, and the enzyme is therefore an attractive target for antiviral action. We have found that the viral protease is inhibited by approximately stoichiometric concentrations of copper or mercury ions. Inactivation by Cu2+ was rapid and not reversed by subsequent exposure to EDTA or dithiothreitol. Direct inhibition by Cu2+ required the presence of cysteine residue(s) in the protease. Thus, a synthetic protease lacking cysteine residues was not inhibited by exposure to copper. However, addition of dithiothreitol as

an

exogenous thiol rendered even the synthetic protease susceptible to inactivation by copper. Oxygen was not required for inactivation of either the wild-type or the synthetic protease. These results provide the basis for the design of novel types of protease inhibitors.

ACCESSION NUMBER:

1999030124 MEDLINE

DOCUMENT NUMBER:

99030124 PubMed ID: 9814959

TITLE:

Chemical inactivation of retroviral infectivity by

targeting nucleocapsid protein zinc fingers: a candidate

SIV vaccine.

AUTHOR:

Arthur L O; Bess J W Jr; Chertova E N; Rossio J L; Esser M

T; Benveniste R E; Henderson L E; Lifson J D

CORPORATE SOURCE:

AIDS Vaccine Program, SAIC/Frederick, National Cancer Institute-Frederick Cancer Research and Development

Center,

Maryland 21702-1201, USA.

CONTRACT NUMBER:

NO1-CO-56000 (NCI)

SOURCE:

AIDS RESEARCH AND HUMAN RETROVIRUSES, (1998 Oct) 14 Suppl

S311-9.

Journal code: ART; 8709376. ISSN: 0889-2229.

PUB. COUNTRY:

United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

199901

ENTRY DATE:

Entered STN: 19990128

Last Updated on STN: 19990128 Entered Medline: 19990111

Although most viral vaccines used in humans have been composed of live AB attenuated viruses or whole killed viral particles, the latter approach has received little attention in research on experimental primate immunodeficiency virus vaccines. Inactivation procedures involving heat

or

formalin appear to adversely affect the viral envelope proteins. Recently we have inactivated human immunodeficiency virus type 1 (HIV-1) with the compound 2,2'-dithiodipyridine (Aldrithiol-2, Aldrich, Milwaukee, WI), which inactivates infectivity of retroviruses by covalently modifying the nucleocapsid zinc finger motifs. HIV-1 inactivated with Aldrithiol-2 retained the conformational and functional integrity of the viral and virion-associated cellular proteins on the viral membrane. We have extended our studies of zinc finger targeted inactivation to simian immunodeficiency virus (SIV) and evaluated

the feasibility of applying the procedures to large scale (>30 1) production and purification of the primate immunodeficiency viruses. There

was no detectable residual infectivity of SIV after treatment with 1 mM Aldrithiol-2 (>5 logs inactivation). Treatment with Aldrithiol-2 resulted in extensive reaction with the nucleocapsid protein of treated virus, as shown by immunoblot and high-performance liquid chromatography (HPLC) analysis. As expected, the virion gp120SU appeared to be completely unreactive with Aldrithiol-2. Sucrose gradient purification and concentration procedures resulted in little

loss

of viral infectivity or virion-associated gp120SU. When tested in a gp120-CD4 dependent cell binding assay, the inactivated virus bound to cells comparably to the untreated virus. Analysis of gp120-CD4 mediated postbinding fusion events showed that the inactivated virus could induce CD4-dependent fusion with efficiencies similar to the untreated virus. Inactivation and processing of primate immunodeficiency viruses by

methods

described here results in highly concentrated virus preparations that retain their envelope proteins in a native configuration. These inactivated virus preparations should be useful in whole killed-particle vaccine experiments as well as laboratory reagents to prepare antisera, including monoclonal antibodies, and to study noninfective virion-cell interactions.